Solution NMR Structure of the NlpC/P60 Domain of Lipoprotein Spr from *Escherichia coli*: Structural Evidence for a Novel Cysteine Peptidase Catalytic Triad

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ABSTRACT: *Escherichia coli* Spr is a membrane-anchored cell wall hydrolase. The solution NMR structure of the C-terminal NlpC/P60 domain of *E. coli* Spr described here reveals that the protein adopts a papain-like α+β fold and identifies a substrate-binding cleft featuring several highly conserved residues. The active site features a novel Cys-His-His catalytic triad that appears to be a unique structural signature of this cysteine peptidase family. Moreover, the relative orientation of these catalytic residues is similar to that observed in the analogous Ser-His-His triad, a variant of the classic Ser-Asp charge relay system, suggesting the convergent evolution of a catalytic mechanism in quite distinct peptidase families.

Bacterial lipoproteins make up a diverse class of membrane-associated proteins that play important roles in a wide range of biological processes and in bacterial pathogenesis (1). In general, lipoproteins are comprised of an N-terminal signal peptide sequence that is cleaved within a conserved lipobox at an invariant cysteine, which in turn is covalently anchored to the bacterial membrane via a diacylglycerol thioether linkage (1–3). The *spr* gene of *Escherichia coli* encodes a 188-residue precursor of a predicted periplasmic surface outer membrane lipoprotein (SWISS-PROT entry SPR_ECOLI; NESG target entry ER541), consisting of a 26-residue N-terminal signal peptidase II recognition sequence culminating in the membrane anchoring cysteine. Its C-terminal domain is a member of the large NlpC/P60 protein domain family (4) (Pfam entry PF00877; MEROPS peptidase clan CA, family C40) comprised of more than 1300 sequences (Pfam 22.0) predominantly from bacteria and featuring both secreted and predicted membrane-bound lipoproteins. While the exact substrate specificity of *E. coli* Spr has not yet been established, the C40 peptidase/NlpC/P60 protein domain families are classified as γ-d-glutamyl-l-diamino acid-endopeptidases which hydrolyze specific peptide linkages in bacterial cell walls and are intimately involved in cell wall hydrolysis during cell growth and division or cell lysis/invasion (4, 5). Accordingly, members of this class of peptidases are potential targets for antibiotic drug discovery.

Here, we present the solution NMR structure of the C-terminal 126-residue NlpC/P60 domain of *E. coli* Spr, Spr[37–162] (molecular mass of 14.4 kDa, pl 10.2), which corresponds to residues 63–188 in the Spr precursor (Figure 1A). The protein adopts a fold whose scaffold is common to the papain superfamily of cysteine peptidases (4). The highly conserved active site of the protein reveals a novel Cys-His-His catalytic triad that appears to be unique to the NlpC/P60 family of enzymes, present in most members of this domain family, including the unpublished structures of two homologues from cyanobacteria, *Nostoc punctiforme* protein 53686717 and *Anabaena variabilis* Q3M7N3 (Figure 1A). To the best of our knowledge, this report represents the first structural description of an active site Cys-His-His catalytic triad in a peptidase, and the first detailed experimental structural perspective on this large and biologically important protein domain family.

The solution NMR structure determination of *E. coli* Spr[37–162] (PDB entry 2K1G; BMRB entry 15603), including the cloning, expression, and purification of 13C- and 15N-enriched protein samples, was performed following standard protocols of the Northeast Structural Genomics Consortium (6) (see the Supporting Information for a complete description of the methods used in this work, as well as a summary of the NMR data and structural statistics for this study). The protein is monomeric in solution, based on gel filtration chromatography, static light scattering, and 15N relaxation data. The structure of *E. coli* Spr[37–162]...
adopts a papain-like α+β fold comprised of four α-helices and a sheet of six antiparallel β-strands arranged in an αααββββααβ topology (Figure 1B). As predicted for the entire NlpC/P60 domain family (4), a highly conserved catalytic cysteine (C68) occurs at the end of a helix (α2) and is packed against a β-sheet core featuring a conserved histidine (H119) from a β-strand (β3). A Consurf (7) analysis of surface features conserved across the NlpC/P60 family demonstrates that highly conserved residues are clustered in a large groove, clearly identifying the active site and substrate binding site of this class of enzymes (Figure 1C). The groove is lined with numerous conserved polar and charged residues, culminating with the catalytic C68 and partially buried H119. A hydrogen bonding network stabilizing neutral side chains revealed by NMR data. In this arrangement, H119 can act as a general base in catalysis, with the second histidine in the triad, H131, serving to properly orient the side chain in the triad (control). pKa values for each are given. (F) Superposition of the active site residues of E. coli Spr[37–162] (PDB entry 2K1G; red), N. punctiforme protein 53686717 (PDB entry 2EVR; green), and human cytomegalovirus serine peptidase (PDB entry 1WPO; blue) (18). The key side chain atoms involved in these catalytic triads are labeled. For the superposition, the Cα and Sα atoms of C68, the Nα atom of H119, and the Nα atom of H131 in the E. coli Spr[37–162] structure were superimposed with the equivalent atoms in the other structures. Hydrogen bond distances: S−Nα, 3.83 ± 0.46 Å for 2K1G, 3.60 Å for 2EVR; N2−Nα, 2.81 ± 0.05 Å for 2K1G, 2.71 Å for 2EVR; O2−N3, 3.07 Å for 1WPO; N3−N2, 3.09 Å for 1WPO. All structure figures were made using PyMol 1.1 (http://www.pymol.org).

Although this function in peptidases is often performed by backbone and side chain (Asn/Gln) amide moieties, there is precedent in the literature for stabilization of the oxyanion intermediate by a tyrosine OH group in certain serine peptidases, including prolyl oligopeptidase (9). We propose that the active catalytic network in this enzyme comprises the thiol group of C68, Nα of H119, Hε2 of H119, and Nα of H131, with both histidines adopting the N2H tautomeric state revealed by NMR data. In this arrangement, H119 can act as a general base in catalysis, with the second histidine in the triad, H131, serving to properly orient the side chain of H119. A hydrogen bonding network stabilizing neutral (imidazole base) histidines, as shown in Figure 1D, would be expected to exhibit reduced histidine pKa values. Indeed, pH titrations of E. coli Spr[37–162] monitored by NMR reveal pKa values of 5.5 for both active site histidines (Figure 1E), compared with a pKa of 6.7 for the histidines in the unstructured hexa-His tag, confirming our picture of this novel catalytic triad at neutral pH.

The structure of the NlpC/P60 domain of E. coli Spr presented here is quite similar to structures of homologous domains in two cell wall hydrolases from cyanobacteria, namely, N. punctiforme protein 53686717 (PDB entry 2EVR) and A. variabilis Q3M7N3 (PDB entry 2HBW), which are
81% identical in sequence to each other. Structural alignments by Dali (10) reveal significant structural similarity between the NMR structure of *E. coli* Spr[37–162] and these two crystal structures (Dali Z scores of 15.3 for 2EVR and 14.8 for 2HBW; Cα rmsds of 2.3 Å for 2EVR and 2.3 Å for 2HBW; sequence identities of 27% for 2EVR and 28% for 2HBW). These 234-residue proteins feature a separate N-terminal SH3-like β-barrel, followed by an NlpC/P60 domain which superimposes very well with the structure of *E. coli* Spr[37–162], in spite of the relatively low level of sequence identity between these bacterial proteins from two distinct phyla. Moreover, the relative orientations of the residues in the catalytic triad are practically identical in the three structures.

The *E. coli* Spr[37–162] structure is also distantly related to two CHAP (cysteine, histidine-dependent amidohydrolases/peptidases) domain family (11) (PF05257; MEROPS clan CA, family C51) cysteine peptidases both featuring a Cys-His-Glu catalytic triad, namely, the N-terminal amidase domain from *E. coli* glutathionylspermidine synthetase (PDB entry 21OB) (12) and a secretory antigen from *Staphylococcus saprophyticus* (PDB entry 2K3A) (13) (Dali Z scores of 7.3 for 21OB and 5.8 for 2K3A; Cα rmsds of 2.9 Å for 21OB and 5.1 Å for 2K3A; sequence identities of 13% for 21OB and 9% for 2K3A). In terms of modeling leverage, defined elsewhere (14), the *E. coli* Spr[37–162] structure has a total modeling leverage value of 482 structural models, and a novel leverage value of 30 models (based on UniProt release 12.8; PSI Blast E < 10⁻¹⁰).

In conclusion, we have described the solution NMR structure of the NlpC/P60 domain of *E. coli* Spr and have identified a novel Cys-His-His catalytic triad in the active site of this cysteine peptidase. To the best of our knowledge, this is the first literature example of this constellation of Cys-His-His active site residues in cysteine peptidases. While β-ketoacyl-acyl carrier protein (ACP) synthases, so-called CHH enzymes, also feature one cysteine and two histidine active site residues (15, 16), the overall protein fold and topology as well as the geometric distribution of the histidines about the cysteine (both point toward the nucleophile) are quite different from those found in the NlpC/P60 domain family (not shown). Interestingly, the analogous rare Ser-His-His catalytic triad, a variation of the classic Ser-His-Asp paradigm first discovered in α-chymotrypsin four decades ago (17), exists in the homodimeric human cytomegalovirus serine peptidase (Pfam entry PF00716; MEROPS clan SH, family S21) (18, 19). In spite of the very different overall fold and opposite orientations of the histidine imidazole rings compared to the structure reported here (i.e., both histidines interact with the preceding residue in the triad via N², not N⁰), the geometries of the catalytic residues are remarkably similar in these Cys-His-His and Ser-His-His triads (Figure 1F). Hence, this appears to be an example of convergent evolution of a catalytic mechanism in disparate peptidase clans. Taken together with the structures for its two distantly related homologues from cyanobacteria, our structure provides a framework for future mutagenesis and biochemical studies on the key residues in the proposed active site (in particular, C68, H119, H131, and Y56), to shed further light on the mechanism of action of this large class of biologically important cell wall hydrolases.

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**SUPPORTING INFORMATION AVAILABLE**

Complete experimental methods used in this study, table of NMR and structural statistics (Table S1), NMR sequential connectivity map (Figure S1), static light scattering data (Figure S2), T² relaxation data (Figure S3), stereoview of the final structural ensemble (Figure S4A), electrostatic surface potential of the lowest-energy conformer (Figure S4B), H⁻¹⁵N HMBC spectrum of Spr[37–162] (Figure S5), and superposition of the Spr[37–162] solution structure with its closest structural homologue (Figure S6). This material is available free of charge via the Internet at http://pubs.acs.org.

**REFERENCES**